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# 1 Examining the Thermal Degradation Behaviour of a Series of 2 Cyanate Ester Homopolymers

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## 11 Abstract

12 A series of thermally stable dicyanate monomers, containing different thermally stable structural  
13 units, viz 2,2'-bis (4-cyanatophenyl)propane (DCDPP), bis-4-cyanato-biphenyl (DCBP), bis-4-  
14 cyanato naphthalene (DCN), 3,3'-bis(4-cyanatophenyl) sulphide (DCTDP) and 3,3'-bis (4-  
15 cyanatophenyl) sulphone (DCDPS), is prepared and the identity of the products confirmed by  
16 FT-IR and NMR spectral methods. The corresponding cyanate homopolymers (designated by the  
17 suffix HP) are prepared and their properties evaluated and compared. The composites were  
18 analysed for their thermal stability and thermal degradation kinetics. The series of  
19 homopolymers exhibit excellent thermal characteristics e.g. relatively high glass transition  
20 temperatures of at least 215 °C, which were inversely proportional to the molecular weight  
21 between the crosslinks, high thermal decomposition temperature, high integral procedural  
22 decomposition temperature (IPDT), and high activation energies for the decomposition of the  
23 cured resins. Determination of their limiting oxygen indices indicates that all the homopolymers  
24 are characterized as 'self-extinguishing' materials.

25 **KEY WORDS:** Cyanate esters, Homopolymers, Curing, Flame retardance, Thermogravimetric  
26 analysis

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28

## 29 **1. INTRODUCTION**

30 Cyanate ester resins have stimulated substantial interest, due to their exclusive combination of  
31 properties, such as low water absorption, low dielectric constant and heat release rate, superior  
32 strength, excellent bonding towards metals, glass and carbon matrices, low volatility while  
33 curing, and high resistance towards high heat and high humid environments. Owing to their  
34 excellent final cured properties, they find application as structural adhesives for making high  
35 temperature resistant and light-weight advanced composites [1,2]. Cyanate esters have definite  
36 advantages over bismaleimide (BMI) resins due to lower typical crosslink densities and higher  
37 flexibility because of the high percentage of oxygen linkages present [3]. These attributes of  
38 cyanate esters are reflected in the higher fracture toughness observed when incorporated into  
39 epoxy resins in comparison with BMIs [4].

40 Cyanate ester resins are primarily used in the field of aerospace materials, in dielectric  
41 components, printed circuit boards, coatings and other applications requiring high temperature  
42 resistant and moisture resistant materials. These applications are the consequence of their high  
43 mechanical strength, high moisture resistance, low dielectric loss, low volatility during the cure,  
44 and low toxicity [5-6]. In recent years, many new cyanate monomers have emerged especially  
45 the dicyanates containing aromatic ether [7], ketone [8,9], 2,7-dihydroxynaphthalene [10],  
46 polyurethane [11], thiophenols [12,13], silicones [14,15] and phosphorus [16,17]. Though  
47 cyanate esters are known to exhibit excellent thermal properties, investigations into the thermal  
48 behaviour, thermal stability and thermal degradation kinetics studies using different  
49 mathematical models under non-isothermal and isothermal conditions are more limited beyond  
50 very basic cyanate esters [18,19]. The trend for increased use of cyanate esters is due to the

51 growing demand for light-weight, low dielectric loss, high heat resistant structures for  
52 manufacturing military aircraft [4,20-27]. Consequently, in this study, a detailed and systematic  
53 evaluation is analyzed and discussed, especially concerning the thermal properties of the  
54 homopolymers.

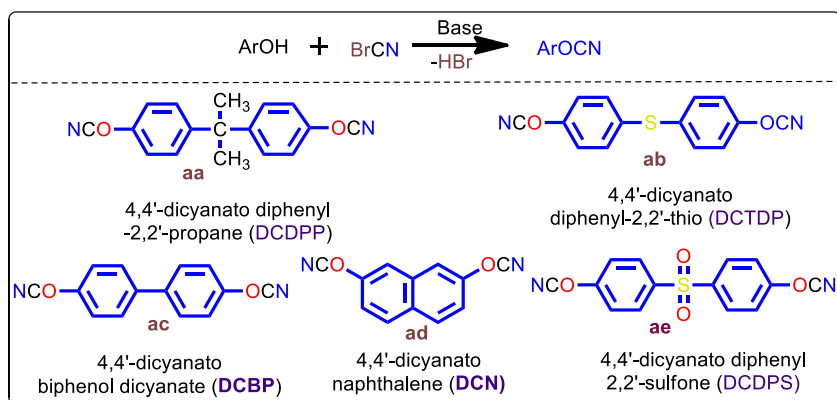
## 55 2. MATERIALS AND METHODS

### 56 2.1. Materials

57 2,2-Bis(4-hydroxyphenyl)propane (99%), 1,4-naphthalene diol (99%), 4,4'-biphenyldiol (97%),  
58 4,4'-thiodiphenol (99%), 4,4'-sulphonyl diphenol (98%), and cyanogen bromide (99%) were  
59 purchased from Aldrich Chemical Company. Triethylamine, acetone and methanol (Analytical  
60 Reagent grade) were purchased from S.D. Fine Chemicals Pvt. Ltd., Mumbai, India.

### 61 2.2. Synthesis of 2,2'-bis(4-cyanatophenyl)propane (DCDPP)

62 A batch scale of cyanate ester (100g) was synthesized at 0 °C by the reaction of cyanogen  
63 bromide (74.6 g, 0.7 mol.) and bisphenol A (81 g, 0.35 mol.). Triethylamine (90 g, 0.89 mol.)  
64 was added to catalyse the reaction and to absorb the evolved HBr to yield salts of tri-  
65 ethylamine hydrobromide (Fig. 1 and Fig. S1). The synthesis was carried out in a three-necked  
66 round-bottomed flask equipped with a mechanical stirrer and a nitrogen inlet was charged a  
67 cooled solution of bisphenol A and cyanogen bromide in acetone. Triethylamine was added  
68 dropwise under continuous stirring in an ice bath and, after complete addition, the reaction  
69 mixture was stirred for a further period of one hour while maintaining the ice bath temperature at  
70 0 °C and filtered under vacuum. The reaction mixture was filtered and, the filtrate was poured  
71 into a large amount of cold distilled water (1L) to precipitate the bisphenol A cyanate ester,  
72 DCDPP, Fig 1(a), from the solution. The crude product was further purified by recrystallization  
73 in methanol: water (1:1 V/V). The product was a white crystalline with 76 g yield (80%) and  
74 m.p. 75-78 °C.



75

76 Fig. 1. Reaction scheme for the preparation of the dicyanate monomers (a-e).

77 The remaining cyanates: DCTDP (RMM 268; m.p. 79 °C solid, white colour), Fig 1(b),  
 78 DCBP (RMM 236; m.p. 82 °C solid, brown colour), Fig 1(c), DCN (RMM 210; m.p. 80 °C,  
 79 solid, dark brown colour), Fig 1(d), and DCDPS (RMM 300; m.p. 81 °C, solid, light brown  
 80 colour), Fig 1(e) were prepared from their respective dihydroxy compounds by employing the  
 81 same procedure [19,20]. All the products were characterized by FT-IR and <sup>13</sup>C NMR  
 82 spectroscopic techniques. Each cyanate ester (100 parts) was homopolymerized (Fig. 2) by  
 83 heating at 140 °C for 3 h, 160 °C for 2 h and followed by a post-curing at 180°C for 4h and 200  
 84 °C for 2 h.

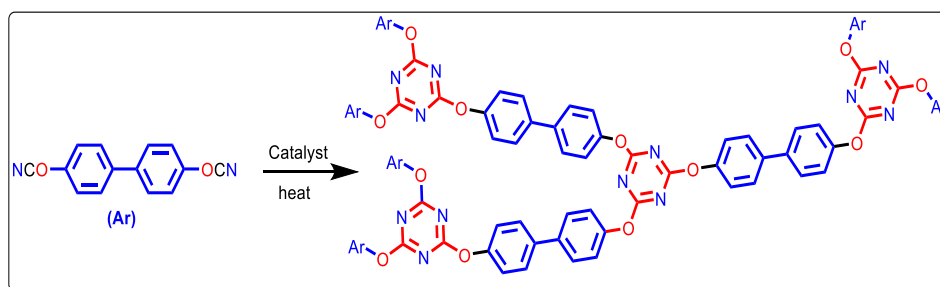
85  
86

Fig. 2. General reaction scheme for the cyclotrimerisation of the dicyanate monomers.

### 87 2.3. Characterisation

88 Fourier Transform Infrared (FT-IR) spectra were obtained using a Nicolet model 20DXB  
89 spectrophotometer with KBr pellets for solid specimens within scanning range of 400-4000  $\text{cm}^{-1}$   
90 at the resolution of  $< 0.1 \text{ cm}^{-1}$ . A JOEL ECA-500 nuclear magnetic resonance (NMR)  
91 spectrometer was used to carry out the analyses at 298K using TMS standard and  $\text{CDCl}_3$  solvent.  
92  $^1\text{H}$  NMR spectra were recorded at 500 MHz and  $^{13}\text{C}$  NMR at 125 MHz. The thermal stabilities of  
93 the cured polymers were determined using TGA Q50-TA thermal analyzer. The  
94 thermogravimetric analysis (TGA) curves were recorded between 30-800  $^\circ\text{C}$  for cured HP  
95 samples (10-15 mg) at a heating rate of  $10^\circ\text{C}/\text{min}$  and under a flowing nitrogen atmosphere (10  
96  $\text{cm}^3/\text{min}$ ). The differential scanning calorimetric (DSC) studies were conducted on DSC Q200  
97 TA instrument at the heating rate of  $5^\circ\text{C}/\text{min}$  between 0-300  $^\circ\text{C}$ . Nitrogen gas flow rate was kept  
98 at the rate of  $10 \text{ cm}^3/\text{min}$ . Scanning electron microscopic (SEM) analysis was performed using a  
99 JEOL 400 microscope on the fractured surface of the cured neat resin applying an accelerating  
100 voltage of 5kV; the fractured samples were first sputtered with carbon.

## 101 3. RESULTS AND DISCUSSIONS

### 102 3.1. Characterization of monomers by FT-IR and $^{13}\text{C}$ NMR spectroscopy

103 FT-IR spectra of the dicyanate monomers (see Supplementary data, Fig. S2) display the  
104 characteristic (O-C $\equiv$ N) doublet around  $2200 \text{ cm}^{-1}$ , confirming the presence of the cyanate group,  
105 while the  $^{13}\text{C}$  NMR spectra (Fig. S3) showed the resonances corresponding to the cyanate  
106 functional groups and all the carbons present in the compounds. The characteristic signals of the  
107 OCN carbons attached to the aromatic rings for DCDPP, DCBP, DCN, DCTDP, and DCDPS  
108 were observed at 116, 109.4, 116.9, 106.5, and 111.6 ppm respectively. The aromatic carbons  
109 appeared in the range 129-188 ppm for DCBP, 118-153 ppm for DCDPP, 118-162 ppm for  
110 DCDPS, 106-154 ppm for DCN, and 116-156 ppm for DCTDP respectively.

## 111 3.2. DSC analysis

### 112 3.2.1. Thermal behaviour

113 From the DSC curves (Fig. S4) of the cured samples of cyanate homopolymers systems,  
114 DCDPP-HP, DCBP-HP, DCN-HP, DCTDP-HP, and DCDPS-HP exhibited their glass transition  
115 temperatures ( $T_g$ ) at 247, 215, 256, 253, and 215 °C respectively. The high  $T_g$  of the  
116 homopolymer systems ( $> 200$  °C) is due to the presence of the triazine rings, formed by the  
117 cyanate monomers. Amongst the homopolymer systems, DCN-HP exhibits the highest  $T_g$ : the  
118 naphthalene moiety has a rigid, planar structure, which packs more readily through  $\pi$ - $\pi$  stacking  
119 [21-22]. The mass loss occurred up to 130 °C is because of the elimination of solvent and  
120 moisture for purification of polymers. Thermal stability of DCDPP-HP shows the highest and  
121 DCN-HP shows lowest due to the fused naphthalene core. Mainly, DCDPP-HP and DCPPS-HP  
122 show higher resistance to heat because of the presence of sulfone and dimethyl propane groups  
123 and which contribute to the single degradation step. On the other hand, the remaining three  
124 homopolymers display two degradation steps: the first due to the single bond scissions and the  
125 second could be the pyrolysis of the cyanurate rings (380 °C- 420 °C). The mass loss occurred up  
126 to 130 °C is because of the elimination of solvent and moisture for purification of polymers.  
127 Thermal stability of DCDPP-HP shows the highest and DCN-HP shows lowest due to the fused  
128 naphthalene core.

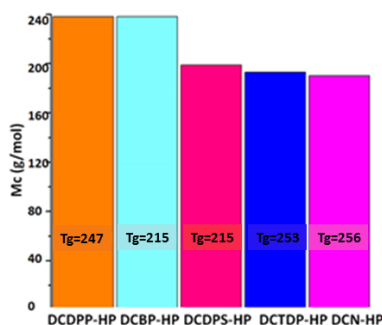
### 129 3.2.2. Estimation of $M_c$

130 The estimation of the molecular weights between adjacent crosslinks ( $M_c$ ) helps to understand  
131 the development of the physical network structure of the polymer since  $M_c$  is inversely  
132 proportional to the crosslink density. The latter is one of the vital structural parameters that aid  
133 knowledge of the influence of changes in the segmental motions, which are reflected in the  
134 mechanical properties of thermoset polymers. When the number of crosslink junctions increases,

135 the crosslink density increases and this to a concomitant increase in  $T_g$ . Hence, the relationship  
 136 between  $T_g$  and  $M_c$  can also be correlated with the crosslink density of the polymer. The  $M_c$   
 137 values for the homopolymer systems presented in this article were estimated using an empirical  
 138 equation 1 [23].

$$139 \quad M_c = \frac{3.9 \times 10^4}{T_g - T_g^0} \quad (1)$$

140  $T_g^0$  is the glass transition temperature of the non-crosslinked polymer.



141

142 Fig. 3. Correlation study presenting molecular weight between the crosslinks ( $M_c$ ) for the  
 143 different cyanate ester homopolymers

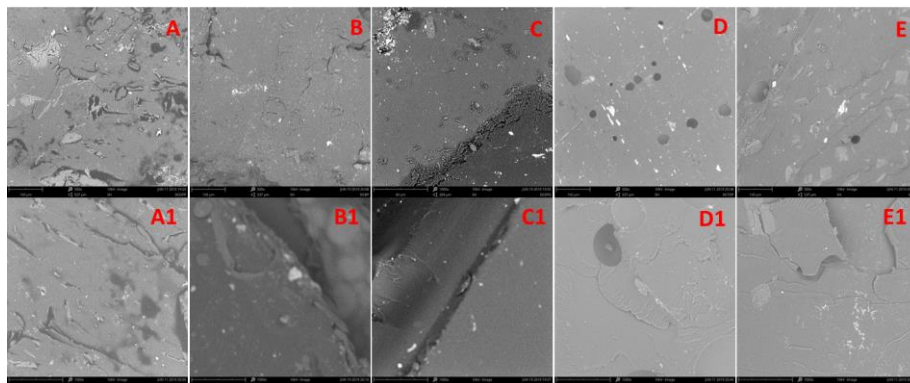
144 Cyanate ester homopolymers show a range of  $M_c$  values from  $198 \text{ g.mol}^{-1}$  to  $240 \text{ g.mol}^{-1}$  (Fig.  
 145 3). DCN-HP has the lowest  $M_c$  value ( $198 \text{ g.mol}^{-1}$ ) following by DCTDP-HP, then DCDPS-HP,  
 146 DCBP-HP, and DCDPP-HP ( $240 \text{ g.mol}^{-1}$ ). Those values depend on the planar structure of the  
 147 aromatic ring and the ability of localized motions of chain segments [24]. In this reason, amongst  
 148 the homopolymer systems, DCN-HP exhibits the lowest  $M_c$  value.

### 149 3.3. Fracture analysis

150 The cured homopolymers were subjected to SEM analysis to analyse the fractured surfaces. In  
 151 general, the surfaces (Fig. 4) show similar, complex morphologies that are typical of shear



152 failure; all the polymers show elastic deformation zones that predominate. Previously, a  
153 commercial cyanate ester (AroCy B-30), which shares an identical chemical structure to DCDPP,  
154 was analysed using similar conditions.[25]



155  
156 Fig. 4. Scanning electron microscopic images of the cured cyanate homopolymers with 500X  
157 (A-E) and 1000X (A1-E1) for DCDPP (A & A1), DCBP (B&B1), DCDPS (C&C1), DCTDP  
158 (D&D1), and DCN (E&E1)

159

### 160 3.4. Thermogravimetric analysis

#### 161 3.4.1. Thermal properties

162 The thermal stabilities of the cured homopolymers were examined using the TGA technique: the  
163 thermal stability of polymers was evaluated by a number of parameters (Table 1) may lead to  
164 contradictory results, in terms of the onset temperature for degradation (DCDPP-HP), lowest rate  
165 of mass loss (DCN-HP), or highest char yield (DCN-HP and DCTDP-HP). The maximum  
166 decomposition temperature (MDT) is the temperature at which the highest rate of thermal  
167 degradation is recorded.

168

169

170

171 Table 1. Thermal properties of cyanate homopolymer

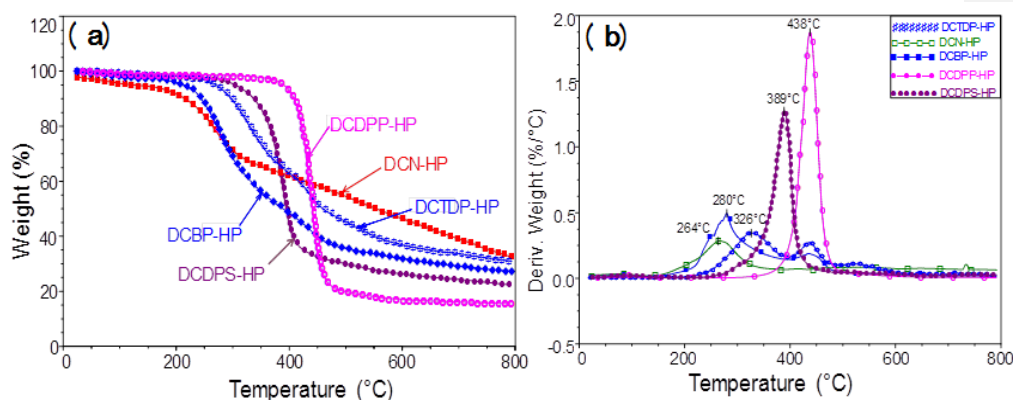
Polymer	Initial decomposition temperature IDT (°C)	Mass loss (%)			Maximum decomposition temperature MDT (°C)	Char residue at 800°C CR (%)
		3	15	30		
DCBP-HP	246	219	273	288	327	28
DCN-HP	229	196	260	274	264	35
DCTDP-HP	250	319	346	441	327	31
DCDPP-HP	388	411	434	446	438	16
DCDPS-HP	323	350	365	377	389	22

172

173 These data were not acquired using hyphenated apparatus in which chemical speciation was  
 174 possible. Consequently, inferences are drawn based on the profile of the thermal degradation and  
 175 comparisons of masses lost, with chemical moieties found within the polymer backbones. The  
 176 TGA data for the homopolymers (Fig. 5a) suggests that, beyond the differences in onset  
 177 temperatures, there are similarities in the degradation mechanism observed for DCBP-HP and  
 178 DCTDP-HP. DCDPS-HP and DCDPP-HP both similarly lose mass in a single drop. The mass  
 179 loss observed up to 150 °C is probably due to the removal of water, which is used for  
 180 recrystallization along with methanol.

181 DCN-HP shows the maximum decomposition temperatures all exceed 260 °C, and this occurs  
 182 around 400 °C for the best performing systems (Fig. 5b), which contain sulphur in the backbone  
 183 structure.

184



185

186 Fig. 5. TGA data (a) and DTG data (b) for the homopolymer systems carried out under N<sub>2</sub>  
 187 atmosphere at a heating rate of 10°C/min.

188 The thermal decomposition of aromatic polycyanurates goes through a common mechanism  
 189 which begins with thermolytic cleavage of the resin backbone and culminates with decyclization  
 190 of the cyanurate rings around 300 °C, followed by char formation; these data are in good  
 191 agreement with previous reports [26-28]. However, the other onsets can be attributed to two  
 192 different species already present in the treated sample and not only formed during TGA  
 193 execution, which showed the first onset around 450 °C corresponding to the decomposition into  
 194 gaseous sulfonyl-di-benzene or propane-2,2-diyldibenzene unit. The char yields observed are  
 195 typical for di-functional aromatic dicyanate homopolymers and DCN-HP, with the highest  
 196 aromatic content, predictably shows the highest char yield (Table 1); DCDPP-HP with the  
 197 greatest aliphatic character, the lowest. The flame-retardant property was found out from the  
 198 limiting oxygen index (LOI) value, using the empirical formulae proposed by Van Krevelen *et*  
 199 *al.* [29]. A numerical index, the LOI represents the minimum concentration of oxygen required  
 200 to support the combustion of a polymer in the particular air mixture. Thus, higher LOI values  
 201 represent better flame retardancy.

202

$$LOI = 17.5 + 0.4CR \quad (2)$$

203 where, *LOI* = limiting oxygen index, and *CR* = yield of char residue at 800°C.

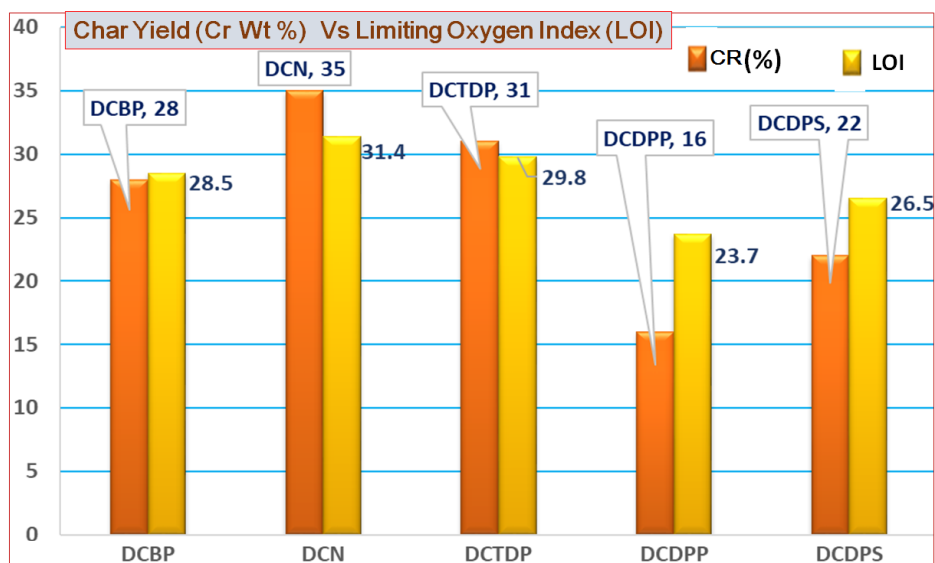
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205 The homopolymer systems show the highest LOI values, which fall between 23-30, where values

206 of LOI &lt; 20.95, LOI &lt; 28.0, and LOI &lt; 100 are considered to represent 'flammable', 'slow-

207 burning', and 'intrinsically non-flammable' materials respectively (Fig. 6).

208

209 Fig. 6. The comparison of char yield vs. LOI to the cyanate system, the labelled (orange is CR  
210 (%), Yellow is LOI (%))

211

212 Using a more detailed description, the polymers possessing LOI  $\geq 20.95$  and  $\geq 26.0$  are

213 considered as 'marginally stable' and 'self-extinguishable' materials, respectively [30-32].

214 According to these criteria, all the homopolymer systems (except DCDPP) have exhibited 'self-

215 extinguishing' characteristics.

216

217 **3.4.2. Determination of activation energy**

218 Thermogravimetric analysis was used for the determination of the kinetics of the thermal  
 219 degradation of the polymers. The thermal degradation of the cured system was carried out at a  
 220 heating rate of 10K min<sup>-1</sup> under a flowing nitrogen atmosphere. The activation energy and order  
 221 of reaction (*n*) were predicted by using the integral methods of the Broido, Horowitz-Metzger,  
 222 and the Coats-Redfern models [334-35] derived from the Arrhenius equation.

223 The basic equation used to describe decomposition reactions is

$$224 \quad \frac{dy}{dt} = k(T)f(y) \quad (34)$$

225 where the rate constant  $k(T)$  and  $f(y)$  were functions of temperature, and conversion respectively  
 226 was defined as

$$227 \quad y = \frac{M_0 - M_t}{M_0 - M_f} \quad (45)$$

228 where  $M_0$ : initial sample weight,  $M_t$  and  $M_f$  were the weight at time  $t$  and final sample weight,  
 229 respectively. Usually,  $k$  is assumed to follow the Arrhenius relationship:

$$230 \quad k = A \exp\left(\frac{-E}{RT}\right) \quad (65)$$

231 The reaction rate may be written as follows.

$$232 \quad \frac{dy}{dt} = \frac{dy}{dT} \frac{dT}{dt} = \beta \frac{dy}{dT} \quad (76)$$

233 Thus, change in mass vs. temperature can be written as

$$234 \quad \frac{dy}{dT} = \frac{A}{\beta} \exp\left(\frac{-E_a}{RT}\right) f(y) \quad (87)$$

235 The Coats-Redfern equation is as follows:

236 The integral form of Eq. 8 from initial temperature,  $T_i$  corresponding to a degree of conversion

237  $m_0$ , to a peak temperature,  $T_{\max}$ , can be written as

$$238 \quad \int_0^y \frac{dy}{f(y)} = \frac{A}{\beta} \int_{T_0}^{T_r} \exp\left(-\frac{E_a}{RT}\right) dT \quad (98)$$

239 Using an approximation, Broido rearranged Eq (87).

$$240 \quad \ln\left[\ln\frac{1}{y}\right] = -\frac{E_a}{R} \frac{1}{T} + \left(\frac{R}{E_a} \frac{A}{\beta} T_{\max}^2\right)_a \quad (109)$$

$$241 \quad \ln\left[\frac{-\ln(1-y)}{T^2}\right] = \ln\frac{AR}{\beta E_a} \left(1 - \frac{2RT}{E_a}\right) - \frac{E_a}{RT} \quad \text{for } n = 1 \quad (110)$$

242 and the modified equation of Horowitz-Metzger is given by:

$$243 \quad \ln(1-y) = \frac{E_a(T-T_p)}{R(T_p)} \quad \text{for } n = 1 \quad (121)$$

244 All of the models used give approximations since the decomposition of the systems involves

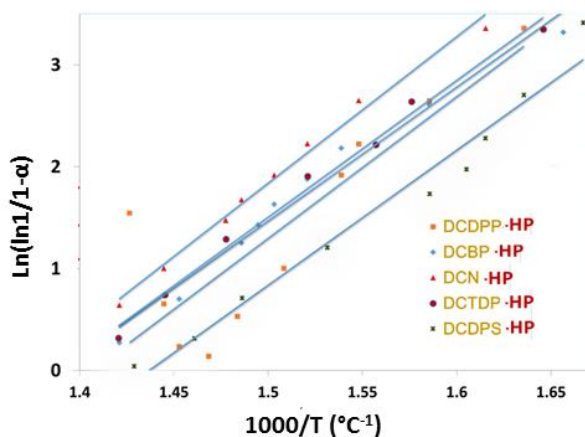
245 several, often coincident or sequential reactions [26-28]. The Horowitz-Metzger model assumes

246 a first-order reaction and uses the simplified exponential integrals to obtain the above equation.

247 The Broido model considers the thermal decomposition process to be a first-order reaction. The

248 Coats-Redfern model, when  $n = 1$ , was considered here for the activation energy calculations

249 [36]. The parameters used were:  $T$  is the absolute temperature,  $\alpha$  is the conversion at temperature  
 250  $T$ ,  $y$  is the fraction of initial molecules and not yet decomposed,  $T_{\max}$  the absolute temperature of  
 251 maximum reaction rate,  $\beta$  is the rate of heating,  $A$  is the frequency factor,  $DT_{\max}$  is the maximum  
 252 decomposition temperature,  $\theta = T - DT_{\max}$ ,  $R$  is the gas constant and  $E_a$  is the activation energy. A  
 253 plot of  $\ln(\ln 1/y)$  in case of Broido's method,  $\ln[-(1-y)/T^2]$  in case of Coats-Redfern method,  
 254 and  $(1-y)$  in the case of the Horowitz-Metzger method; vs.  $1000/T$  for major degradation events  
 255 yielded plots with linear portions. The changes in gradients are consistent with the different steps  
 256 in the thermal degradation mechanism [26]. The cured samples of homopolymer systems were  
 257 subjected to the kinetic analysis and are shown in Fig. 7, for the Horowitz-Metzger model.



258  
 259 Fig. 7. The plots for the calculation of activation energies for homopolymers using the Horowitz-  
 260 Metzger model

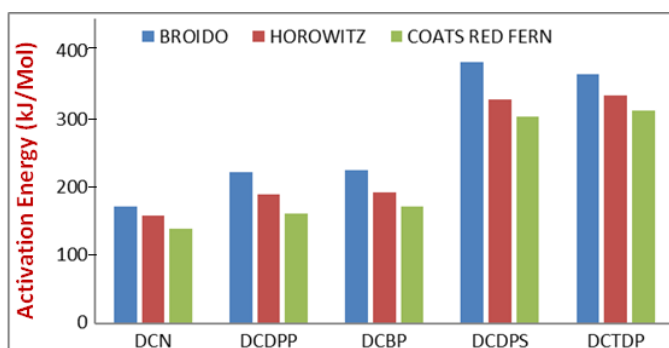
261 The kinetic parameters and the correlation coefficient ( $R^2$ ) values of each system are summarized  
 262 in Table 2.

263 Table 2 Comparison of kinetic parameters for the thermal degradation of the homopolymers using  
 264 different models.

Polymer	Models					
	Broido		Horowitz-Metzger		Coats-Redfern	
	$E_a$ (kJ.mol <sup>-1</sup> )	$R^2$	$E_a$ (kJ.mol <sup>-1</sup> )	$R^2$	$E_a$ (kJ.mol <sup>-1</sup> )	$R^2$
DCBP-HP	225	0.952	193	0.935	173	0.919
DCN-HP	173	0.970	159	0.964	140	0.953
DCTDP-HP	367	0.959	336	0.950	314	0.943
DCDPP-HP	223	0.933	190	0.899	163	0.866
DCDPS-HP	385	0.995	329	0.993	305	0.991

265  $E_a$  = activation energy (kJ.mol<sup>-1</sup>);  $R^2$  = correlation coefficient.

266 While there are some differences in the activation energies ( $E_a$ ) calculated, the trends observed  
 267 are consistent between the different models and the  $E_a$  values were generally found to be in the  
 268 following order for the models applied: Broido>Horowitz-Metzger>Coats-Redfern (Fig. 8). In  
 269 the homopolymers, the  $E_a$  values derived using the Broido model fell between 173 - 385kJ/mol.



270  
 271 Fig. 8. Comparison of activation energies for the thermal degradation of the homopolymers using  
 272 different models.

273  
 274 These values obtained for all the neat resins are higher than values that have previously been  
 275 reported for the bisphenol E cyanate ester based on bisphenol E, 2,2'-bis(4-  
 276 cyanatophenyl)ethylidene, (67 kJ mol<sup>-1</sup>) [37] and a cyanate ester functional benzoxazine (100 kJ



277 mol<sup>-1</sup>) [38] respectively. Within the data set, the highest activation energies calculated for the  
278 thermal degradation of the homopolymers were obtained for the sulphur-containing  
279 functionalized cyanate monomers (DCTP and DCDPS). The presence of sulphur plays a vital  
280 role and the evolution of nonflammable gases (*e.g.* oxides of sulphur) during the degradation  
281 may condense on the remaining polymer, thus diluting oxygen concentration at the polymer  
282 surface, starving the flame, and serving as a free-radical flame front scavenger to inhibit  
283 degradation of the polymers [39].

284

285

286 Five polycyanurate homopolymers displaying similar degrees of crosslink density, but  
287 differing in terms of their molecular rigidity, were analysed for their thermal degradation  
288 behavior, flame retardancy, and fracture properties. The homopolymers are all based on aromatic  
289 monomers and so the high carbon content in the structure of homopolymers and presence of  
290 nitrogen yields moderately high char yields (ranging from 16-35%, with DCDPP, containing an  
291 aliphatic bridge, having the lowest value, and DCN, with no aliphatic character, the highest). The  
292 activation energies (determined using the Broido model) for the decomposition behaviour of the  
293 homopolymers revealed a wide variation, from 173 kJ/mol for DCN to 385 kJ/mol for DCDPS,  
294 with the highest values in the data set being observed for homopolymers derived from simple  
295 monomers.

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